

Research Article

Morpho-phylogenetic evidence reveals *Pseudolomaantha thailandica* gen. et sp. nov. and *Submultiguttulispora multiseptata* gen. et sp. nov. in Chaetosphaeriaceae

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Abstract

Anamorphic chaetosphaeriaceous fungi exhibit high morphological variability and are distributed worldwide across terrestrial and aquatic habitats. During an ongoing taxonomic study of microfungi, two intriguing chaetosphaeriaceous hyphomycetes were collected from dead wood and dead bamboo stems in China and Thailand. A polyphasic approach, combining morphological characteristics and phylogenetic analysis of LSU and ITS sequence data, revealed that these fungi represent two new genera within Chaetosphaeriaceae. Pseudolomaantha and Submultiguttulispora are proposed for these new genera, and they exhibit non-phialidic and phialidic asexual morphs, respectively. Pseudolomaantha thailandica gen. et sp. nov. is characterized by a sporidesmium-like asexual morph with macronematous, mononematous conidiophores; monoblastic conidiogenous cells, and pyriform to obclavate, rostrate conidia bearing an apical appendage. Submultiguttulispora multiseptata gen. et sp. nov. is distinguished by macronematous, mononematous conidiophores, mono- to polyphialidic conidiogenous cells, and fusiform or ellipsoidal-fusiform, pale brown to olive green to brown conidia with filiform, hyaline appendages at both ends. Detailed descriptions, illustrations, and notes on the new collections are provided, along with a key to non-phialidic hyphomycetous genera in Chaetosphaeriaceae.

Key words: 4 new taxa, asexual morph, Sordariomycetes, sporidesmium-like fungus, taxonomy

Introduction

Chaetosphaeriales was established by Huhndorf et al. (2004) to accommodate the family Chaetosphaeriaceae based on morphological characteristics and phylogenetic analysis of LSU sequence data. Currently, four families, *viz.*, Chaetosphaeriaceae, Helminthosphaeriaceae, Leptosporellaceae, and

Linocarpaceae are recognized within this order (Hyde et al. 2020; Wijayawardene et al. 2022). The estimated stem age of Chaetosphaeriales is approximately 158 MYA, based on divergence time analysis (Hyde et al. 2020).

Chaetosphaeriaceae was invalidly introduced by Locquin (1984) without a formal description but was later validly re-established by Réblová et al. (1999) to accommodate Chaetosphaeria and its relatives. Since its re-establishment, the family has shown remarkable diversity with a significant increase in the number of genera and species described in recent years (Lin et al. 2019; Yang et al. 2019; Réblová et al. 2020, 2021a, b, c, d, e, 2022, 2024; Wijayawardene et al. 2022; Wu and Diao 2022, 2023; Réblová and Nekvindová 2023; Yang et al. 2023; Hyde et al. 2024a). Wu and Diao (2022) conducted a comprehensive study of the anamorphic Chaetosphaeriaceae, which included 89 genera, establishing the family as one of the largest within Sordariomycetes (Wijayawardene et al. 2022). Their study also provided identification keys for most genera (Wu and Diao 2022). Following this, Réblová et al. (2022) and Réblová and Nekvindová (2023) carried out systematic reviews of chloridium-like morphotypes, resulting in the addition of seven new genera to Chaetosphaeriaceae. In subsequent years, five more new genera, viz., Gongromerizella, Neocirrenalia, Paragongromeriza, Pseudophialocephala and Pseudostriatosphaeria, were introduced based on morphology and phylogeny (Manawasinghe et al. 2022; Yang et al. 2023; Wu and Diao 2023; Tian et al. 2024; Zhang et al. 2024). Réblová et al. (2024) re-evaluated species in genera Exserticlava, Phaeostalagmus, Phialocephala, and several chalara- and stanjehughesia-like fungi, which led to the establishment of three new genera in Chaetosphaeriaceae. On the other hand, Ellisembia was removed from Chaetosphaeriaceae and reclassified under Sporidesmiaceae (Delgado et al. 2024; Hyde et al. 2024a). Hyde et al. (2024a) accepted 107 genera in Chaetosphaeriaceae.

The sexual morph of Chaetosphaeriaceae is characterized by perithecial, papillate, globose to subglobose, setose, dark brown to black ascomata; unitunicate, clavate to cylindrical asci with a J-, apical ring; and 0-3-septate, fusiform, cylindrical to ellipsoid, hyaline to brown ascospores, often with guttules, a sheath, or appendages (Réblová et al. 1999; Réblová and Gams 2000; Hyde et al. 2020; Wu and Diao 2022). The asexual morphs of Chaetosphaeriaceae include both hyphomycetes and coelomycetes. Coelomycetous morphs are characterized by stromatic, cupuliform or globose, unilocular, setose conidiomata; numerous, septate, ovoid to cylindrical setae; 4-6-septate, unbranched, pigmented conidiophores; integrated, holoblastic or enteroblastic, phialidic conidiogenous cells with conspicuous periclinal thickening at an attenuated apex; and aseptate, hyaline to brown conidia with tubular appendages at the ends (Hashimoto et al. 2015; Hyde et al. 2020; Li et al. 2020). Hyphomycetous morphs are further divided into phialidic and non-phialidic anamorphs. Phialidic anamorphs exhibit macronematous, mononematous, septate, pigmented conidiophores; mono- or polyphialidic conidiogenous cells that proliferate percurrently or sympodially, often with funnel-shaped collarettes; and aggregated, fusiform, allantoid, cylindrical or doliiform conidia, which are mostly hyaline but sometimes pigmented, and often possess filiform appendages (Réblová 2004; Fernández and Huhndorf 2005; Liu et al. 2016; Lin et al. 2019; Luo et al. 2019; Yang et al. 2019; Réblová et al. 2020, 2021a, b, d). Non-phialidic fungi in Chaetosphaeriaceae are predominantly characterized by sporidesmium-like asexual morphs (Ellis 1971, 1976; Wu and Zhuang 2005; Wu and Diao 2022; Yang et al. 2023; Delgado et al. 2024). These taxa are primarily saprobic, occurring on various plant substrates in both terrestrial and aquatic habitats, with some species also found in soil or as fungicolous taxa (Hughes and Kendrick 1968; Perera et al. 2016; Hyde et al. 2018; Réblová et al. 2020, 2021d; Wu and Diao 2022; Zhang et al. 2022; Calabon et al. 2023; Yang et al. 2024; Zhang et al. 2024).

In this study, we aim to introduce two new genera, *Pseudolomaantha* and *Submultiguttulispora*, to accommodate two new species, *P. thailandica* and *S. multiseptata*, respectively. Evidence from morphology and phylogenetic analysis of a combined LSU and ITS sequence dataset supports the establishment of these two new genera (*Pseudolomaantha* and *Submultiguttulispora*) within Chaetosphaeriaceae, Chaetosphaeriales, Sordariomycetes.

Material and methods

Collections, isolation and conservation

Samples of dead bamboo stems and wood were collected from Thailand and China. The collection information of the samples was noted (Rathnayaka et al. 2024), and the samples were taken to the laboratory in zip-lock plastic bags and subsequently examined using the methods described in Senanayake et al. (2020). Morphological observations of the fungal colonies on natural substrates were conducted using a stereomicroscope (Leica EZ4 Microsystems (Schweiz) AG, Singapore). A detailed examination of fungal structures was carried out using a Nikon ECLIPSE Ni compound microscope (Nikon, Japan) and photographed with a Nikon DS-Ri2 digital camera attached to the microscope. Measurements of fungal structures were made using Tarosoft® Image Frame Work, and images used in figures were processed and assembled with Adobe Illustrator CS6 (Adobe Systems, San Jose, CA, USA).

Single-spore isolations were performed on water agar (WA), and germinated spores were transferred to potato dextrose agar (PDA) to obtain pure cultures (Chomnunti et al. 2014). Dried specimens were deposited in the Herbarium of Mae Fah Luang University (MFLU), Chiang Rai, Thailand, and the Herbarium of Cryptogams, Kunming Institute of Botany, Academia Sinica (HKAS), Kunming, China, and the Herbarium of Guizhou Academy of Agricultural Sciences (GZAAS), Guiyang, China. Pure cultures were deposited in the Mae Fah Luang University Culture Collection (MFLUCC) and the Kunming Institute of Botany Culture Collection (KUNCC). Faces of Fungi and Index Fungorum numbers were registered following the guidelines of Jayasiri et al. (2015) and Index Fungorum (http://www.indexfungorum.org/Names/Names.asp; accessed on 15 November 2024)

DNA extraction, PCR amplification and sequencing

Pure cultures were incubated at 25 °C-28 °C for one month. Fresh fungal mycelia were scraped from the surface of the colonies and transferred to a 1.5 mL microcentrifuge tube using a sterilized scalpel for genomic DNA extraction. Genomic DNA was extracted using the Biospin Fungus Genomic DNA Extraction Kit (Biospin Fungus Genomic DNA Extraction Kit, BioFlux®,

Shanghai, China) following the manufacturer's instructions. The large subunit of ribosomal DNA (LSU) and the internal transcribed spacer (ITS) gene regions were amplified using primers LR0R and LR5 (Vilgalys and Hester 1990) and ITS5 and ITS4 (White et al. 1990), respectively. Polymerase chain reaction (PCR) was performed in a 50 μL reaction mixture containing 2 μL of DNA template, 2 μL of each forward and reverse primer (10 μM), 25 μL of 2 × Taq PCR Master Mix with blue dye (Sangon Biotech, China), and 19 μL of distilled—deionized water. Amplification conditions for the LSU and ITS regions followed the protocol described by Zhang et al. (2022). The quality of PCR products was assessed using 1% agarose gel electrophoresis stained with ethidium bromide. Purification and sequencing of PCR products were performed by Beijing Qingke Biotechnology Co., Ltd.

Phylogenetic analyses

Original sequences were verified using BioEdit v. 7.1.3.0 (Hall 1999), and were assembled using SeqMan v. 7.0.0 (DNASTAR, Madison, WI, USA). The newly generated sequences were subjected to BLAST searches in GenBank to determine closely related taxa. Taxa used in the phylogenetic analysis for Chaetosphaeriaceae were selected and obtained from previous studies and GenBank (Wu and Diao 2022; Zhang et al. 2022; Réblová and Nekvindová 2023; Réblová et al. 2024). Sequence alignments for each locus were aligned using the online multiple alignment program MAFFT v.7 (http://mafft.cbrc.jp/alignment/server/, accessed September 2024; Katoh et al. 2019). The alignments were visually checked and manually improved where necessary using BioEdit v. 7.1.3.0 (Hall 1999). LSU and ITS sequences were combined using SequenceMatrix 1.7.8 (Vaidya et al. 2011). Sequences generated in this study were deposited in GenBank (Table 1).

The fasta files were converted to formats required for the AliView program (Larsson 2014), PHYLIP for maximum likelihood analysis (ML), and NEXUS for Bayesian inference (BI). Phylogenetic analyses were performed through the CIPRES science Gateway CIPRES science Gateway V. 3.3 (https://www.phylo. org/portal2/home.action; Miller et al. 2010). Maximum likelihood analysis was performed using RAxML-HPC v.8 tool with rapid bootstrap analysis, followed by 1000 bootstrap replicates (Miller et al. 2010; Stamatakis 2014). The final tree was selected from the suboptimal trees of each run by comparing likelihood scores under the GTRGAMMA substitution model. Bayesian analysis was performed in MrBayes 3.2.7a (Ronquist et al. 2012). The best-fit substitution model GRT + I + G was decided for all two genes by MrModeltest 2.3 under the Akaike Information Criterion (AIC) (Nylander 2004). The Markov Chain Monte Carlo (MCMC) sampling approach was used to calculate posterior probabilities (PP) (Rannala and Yang 1996; Huelsenbeck 2001; Zhaxybayeva and Gogarten 2002). Four simultaneous Markov chains were run for 1 million generations, with trees sampled every 100 generations, resulting in 10,000 trees. The first 2,000 trees, representing the burn-in phase of the analyses, were discarded, and the remaining trees were used for calculating posterior probabilities (PPs) in the majority rule consensus tree (Larget and Simon 1999).

Phylogenetic trees were visualized using FigTree v.1.4.4 (Rambaut 2014), and the layouts were reorganized using the methods described by Xie et al.

Table 1. Chaetosphaeriaceae taxa used in the phylogenetic analysis, and their corresponding GenBank accession numbers.

Taxon	Strain	Status	ITS	LSU
Achrochaeta rivulata	CBS 148186		OR286508	OR286551
Achrochaeta talbotii	ICMP 15161		MT454480	MT454495
Aciculadictyochaeta luquillensis	SMH 2973		N/A	AF466074
Adautomilanezia caesalpiniae	CC-LAMIC 102/12	T	KX821777	KU170671
Anacacumisporium appendiculatum	HMAS 245593	T	KP347129	KT001553
Anacraspedodidymum submerum	YMF1.4176	T	MK165445	MK165443
Arcuatospora novae-zelandiae	CBS 109474		MW984569	MW98455
Arcuatospora seorsa	CBS 147510	Т	MW984572	MW98455
Aunstrupia nodipes	NN043149		OL627566	OL655011
Brachydictyochaeta antillana	NN058987		OL627951	OL655147
Brachydictyochaeta bulliformis	NN076027		OL628023	OL655155
Brunneodinemasporium brasiliense	CBS 112007	Т	JQ889272	JQ889288
Brunneodinemasporium jonesii	GZCC 16-0050	T	KY026058	KY026055
Cacumisporium acutatum	CBS 101312	•	AF178553	AF178553
Cacumisporium acutatum	CBS 101315	Т	OR134682	OR134626
Cacumisporium capitulatum	CBS 101313	1	OR134683	OR134627
Caliciastrum bicolor	ICMP 15136	Т	OR134689	OR134627
Caliciastrum bicolor	PRA-21507	T	N/A	OR134634
	CBS 734.83	T	OR134691	OR134636
Caligospora dilabens				
Caligospora dilabens	CBS 735.83	T	MH861684	N/A
Caligospora pannosa	CBS 551.89	T	OR134692	OR134637
Calvolachnella guaviyunis	CBS 134695	T	KJ834524	KJ834525
Capillisphaeria crustacea	CBS 144665		OR134695	OR134640
Capillisphaeria crustacea	ICMP 15139		OR134696	OR134641
Catenularia angulospora	MFLUCC 18-1331	_	MK828638	MK835840
Catenularia catenulata	DLUCC 0891	T	MK828637	MK835838
Catenularia minor	PRM 900544	Т	MW987827	MW987822
Chaetosphaeria guttulata	MFLUCC 17-1703	T	MK828636	MK835837
Chaetosphaeria innumera	M.R. 3775		OR134699	OR134644
Chaetosphaeria innumera	CBS 145639		OP455358	OP455464
Chaetosphaeria mangrovei	MCD 069	Т	MG813821	MG813820
Chaetosphaeria polygonalis	GZCC 20-0438	Т	OP377861	OP377946
Chalarodes obpyramidata	PDD 119364		MW987828	MW987823
Chloridium bellum	CBS 709.73A	T	OP455360	OP455466
Chloridium caesium	CBS 145633		OP455367	OP455474
Chloridium gamsii	CBS 667.75	T	OP455415	OP455522
Chloridium virescens	CBS 145481		OP455439	OP455547
Codinaea assamica	CBS 139907	T	OL654077	OL654134
Codinaea fertilis	IMI 233824		OL654080	OL654137
Codinaea paniculata	CBS 145098	Т	MT118230	MT118201
Codinaeella lambertiae	CBS 143419	Т	OL654084	OL654141
Codinaeella minuta	CBS 280.59		OL654090	OL654147
Codinaeella parvilobata	CBS 144536	Т	OL654100	OL654157
Conicomyces pseudotransvaalensis	HHUF 29956	Т	LC001710	LC001708
Craspedodidymum elatum	NN042874		OL627547	OL655004
Cryptophiale udagawae	GZCC 18-0047		MN104608	MN104619
Cryptophialoidea fasciculata	MFLU 18-1499		MH758195	MH758208
Curvichaeta curvispora	ICMP 15115	Т	OR134705	OR134650
Curvichaeta curvispora	ICMP 15118		OR134706	OR134651
Dendrophoma cytisporoides	CBS 144107		MT118234	MT118205
Dictyochaeta callimorpha	ICMP 15130		MT454483	MT454498
Dictyochaeta fuegiana	ICMP 15153	Т	MT454487	EF063574

Taxon	Strain	Status	ITS	LSU
Dictyochaeta querna	CBS 145503		MT454489	MT454503
Dinemasporium cruciferum	HHUF 30001		AB900895	AB934039
Dinemasporium pseudoindicum	CBS 127402	T	JQ889277	JQ889293
Ericiosphaeria spinosa	S.M.H. 2754	T	MW984575	AF466079
Eucalyptostroma eucalypti	CBS 142074	Т	KY173408	KY173500
Eucalyptostroma hongluosiense	NN076613		OL628127	OL655185
Eucalyptostromiella beijingensis	NN078016		OL628501	OL655251
Exserticlava vasiformis	TAMA 450		N/A	AB753846
Exserticlavopsis chlorotunicata	S.M.H. 1565	T	N/A	AF466064
Falholtia kaohsiungensis	NCYU108K3-1-1	T	MT939301	MT939304
Falholtia kaohsiungensis	NN050711		OL627699	OL655083
Flectospora laminata	CBS 112964	Т	MW984576	MW984558
Fuscocatenula submersa	MFLUCC 18-1342	T	MK828634	MK835835
Fuscocatenula variegata	NN055332		OL627817	OL655124
Fusichloridium cylindrosporum	CBS 101429	T	OR134709	OR134653
Fusichloridium cylindrosporum	CBS 101430		OR134710	OR134654
Geniculoseta preussii	CBS 263.75		OR134713	OR134657
Geniculoseta preussii	CBS 145478		OR134714	OR134658
Gongromeriza myriocarpa	CBS 264.76		AF178552	AF178552
Gongromeriza myriocarpa	CBS 141.53	Т	OP455456	OP455564
Gongromeriza pygmaea	IMI 506815	1	OR134724	OR134668
Gongromerizella pachytrachela	CBS 645.75	Т	OP455461	OP455569
Gongromerizella pini	CBS 146011	T	MT223787	MT223882
Gongromerizella silvana	CBS 171.76	T	OR134729	OR134673
Infundibulomyces cupulatus	BCC 11929	T	EF113976	EF113979
Infundibulomyces oblongisporus	BCC 13400	T	EF113977	EF113980
Kionochaeta microspora	GZCC 18-0036	T	MN104607	MN104618
Kionochaeta microspora Kionochaeta ramifera	MUCL 39164	ı	MW144421	MW144404
Kionochaetiella ivoriensis	CBS 374.76	Т	MH860988	
				MH872758
Kylindrochaeta lignomollis	S.M.H. 3015	T	EU037896	AF466073
Leptosporella arengae	MFLUCC 15-0330	T	MG272255	MG272246
Leptosporella bambusae	MFLUCC 12-0846	T	KU940134	KU863122
Linkosia multiseptum	CGMCC 3.20786	T	OL627557	OL655008
Linkosia rostrata	CGMCC 3.20790	T	OL627662	OL655059
Lomaantha aquirostrata	GZCC 20-0503	T	OP377802	OP377901
Lomaantha aurantiaca	CBS 126743	T	HM241692	HM241692
Lomaantha aurea	CBS 144403	T	MH836375	MH836376
Lunatochaeta shenzhenensis	CGMCC 3.20757	Т	OL628577	OL655258
Menispora caesia	CBS 145022		OL654107	OL654164
Menispora ciliata	CBS 122131	Т	EU488736	OL654165
Menispora tortuosa	CBS 117553		OL654111	OL654169
Menisporopsis pirozynskii	MUCL 47217		MW984579	MW984561
Menisporopsis theobromae	MUCL 41079		MW984580	MW984562
Morrisiella indica	NN042908		OL627551	OL655005
Morrisiella indica	NN044710		OL627629	OL655037
Multiguttulispora dimorpha	CBS 140002		MW984582	MW984564
Multiguttulispora triseptata	IMI 353690		MW984584	MW984566
Nawawia filiformis	MFLUCC 17-2394		MH758196	MH758209
Neonawawia malaysiana	CPC 16757	Т	GU229886	GU229887
Neopseudolachnella acutispora	MAFF 244358	Т	AB934065	AB934041
Neopseudolachnella magnispora	MAFF 244359	Т	AB934066	AB934042
Neocirrenalia nigrospora	MFLUCC 18-0418		OP377888	OP377974
Nimesporella capillacea	IMI 358908	Т	OL654114	OL654171
Paliphora intermedia	CBS 896.97	ı	MH862682	EF204501
Papillospora hebetiseta	CBS 102340	Т	AF178549	AF178549

Taxon	Strain	Status	ITS	LSU
Paraceratocladiella polysetosa	NN044119		OL627605	OL655027
Paraceratocladium silvestre	NN055375		OL627830	OL655132
Paracryptophiale pirozynskii	CGMCC 3.20706	T	OL627641	OL655047
Paragaeumannomyces panamensis	S.M.H. 3596	T	AY906948	MT118218
Paragaeumannomyces rubicundus	S.M.H. 3221	T	MT118242	MT118224
Phaeodischloridium aquaticum	MFLUCC 18-1341	T	MK828639	MK835841
Phialoarthrobotryum triseptatum	CBS 120.84	T	MH861706	MH873417
Phialogeniculata guadalcanalensis	MFLUCC 18-0260	T	MK828625	MK835825
Phialogeniculata guadalcanalensis	NN044662		OL627622	OL655032
Phialosporostilbe scutiformis	MFLUCC 17-0227	Т	MH758194	MH758207
Phialosporostilbe scutiformis	MFLUCC 22-0053		ON678180	ON678145
Phialoturbella calva	ICMP 23826	Т	MW984585	MW984567
Phialoturbella lunata	MFLUCC 18-0642	Т	MK828624	MK835824
Polynema podocarpi	CBS 144415	Т	MH327797	MH327833
Pseudodinemasporium fabiforme	CBS 140010		KR611889	KR611906
Pseudolachnea fraxini	CBS 113701	Т	JQ889287	JQ889301
Pseudolachnea hispidula	MAFF 244365	-	AB934072	AB934048
Pseudolachnella asymmetrica	MAFF 244366	Т	AB934073	AB934049
Pseudolachnella scolecospora	MAFF 244379		AB934086	AB934062
Pseudolomaantha thailandica	MFLUCC 24-0521	Т	PQ625465	PQ625467
Pseudothozetella lunata	CGMCC 3.20661	T	OL628034	OL655157
Psilobotrys minutus	CBS 877.73	1	OR134733	OR134677
Psilobotrys minutus	CBS 145632		OR134734	OR134678
Rattania setulifera	GUFCC 15501	Т	GU191794	HM171322
	CGMCC 3.20794	T	OL627701	OL655085
Riisgaardia longispora				
Riisgaardia obclavata	CGMCC 3.20787	Т	OL627568	OL655013
Riisgaardia vermiculata	NN042952		OL627555	OL655007
Spadicocephala fusca	CBS 301.85		AF486122	MH873571
Spadicocephala fusca	CBS 300.85		MH861882	MH873570
Spicatispora fennica	CBS 101641	_	OR134735	OR134679
Sporendocladia beijingensis	CGMCC 3.20738	T	OL628290	OL655217
Sporendocladia fumosa	NN047731		OL627669	OL655065
Sporoschisma hemipsilum	MUCL 56487		MW987829	MW987824
Sporoschisma mirabile	CBS 144794		MW987830	MW987825
Stanjehughesia hormiscioides	S.M.H.2794		N/A	AF466060
Stilbochaeta malaysiana	IMI 312436	Т	OL654121	OL654178
Stilbochaeta ramulosetula	IMI 313452	T	OL654124	OL654181
Striatosphaeria castanea	CBS 145352	T	MT118244	MT118229
Striatosphaeria codinaeophora	M.R. 1230		AF178546	AF178546
Submultiguttulispora multiseptata	KUNCC 23-14145	Т	PQ625466	PQ625468
Tainosphaeria cecropiae	CBS 101687	Т	MW984586	MW984568
Tainosphaeria crassiparies	S.M.H. 1934	Т	MW984587	AF466089
Tainosphaeriella aquatica	MFLUCC 17-2370	Т	MZ161197	MZ161195
Tainosphaeriella thailandense	MFLUCC 18-1282	Т	MZ161198	MZ161196
Thozetella cristata	CBS 101112		OL654126	OL654183
Thozetella tocklaiensis	CBS 378.58	Т	OL654128	OL654185
/erhulstia biformis	NN077655		OL628434	OL655237
Verhulstia trisororum	CBS 143234	Т	MG022181	MG022160
Zanclospora novae-zelandiae	ICMP 15781	· T	MW144429	MW14441
Zanclospora ramifera	ICMP 22738	- -	MW144433	MW14441
Zanciospora iberica	CBS 130426	T	KY853480	KY853544
Larrorospora incrioa	ODO 100420	1	111000400	111000044

 $Note: status: T denotes type strains; "N/A" indicates data unavailable in GenBank. The newly generated sequences are indicated in {\bf bold}.$

(2023) and finalized with Adobe Illustrator CS6 software (Adobe Systems, USA). Sequences generated from our collections were deposited in Gen-Bank and are listed in Table 1. Decisions regarding the discovery of new species or records were made following the guidelines of Maharachchikumbura et al. (2021).

Phylogenetic analysis results

The partial LSU-ITS nucleotide sequences were used to determine the phylogenetic position of the new taxa within the family Chaetosphaeriaceae. The concatenated sequence matrix comprises 157 ingroup taxa of Chaetosphaeriaceae and two outgroup taxa, *Leptosporella arengae* (MFLUCC 15–0330) and *L. bambusae* (MFLUCC 12–0846). After alignment, the dataset contained 1,450 characters (LSU: 861 bp, ITS: 589 bp), including 853 distinct alignment patterns, with 11.93% comprising undetermined characters or gaps. Base frequencies were as follows: A = 0.224314, C = 0.274605, G = 0.307808, and T = 0.193272. Substitution rates were AC = 1.327038, AG = 1.998330, AT = 1.575283, CG = 0.648947, CT = 6.385392, and GT = 1.000000, with a tree length of 12.245369. The distribution shape parameter (α) was calculated as 0.317788. The ML and BI trees displayed similar topologies with no significant differences. The best-scoring RAxML tree is shown in Fig. 1, with a final likelihood value of -31034.684968.

Our two isolates were identified as *Pseudolomaantha thailandica* gen. et sp. nov. and *Submultiguttulispora multiseptata* gen. et sp. nov. in Chaetosphaeriaceae. *Pseudolomaantha* shares a sister relationship with a clade comprising *Caliciastrum, Caligospora*, and *Craspedodidymum*, while *Submultiguttulispora* forms a separate clade within Chaetosphaeriaceae that is close to *Multiguttulispora*. Both genera represent distinct, independent lineages and do not belong to any existing genera within Chaetosphaeriaceae.

Taxonomy

Pseudolomaantha J.Y. Zhang, Y.Z. Lu & K.D. Hyde, gen. nov.

Index Fungorum: IF903140 Facesoffungi Number: FoF16983

Etymology. The name refers to the new genus's similarity to the genus "Lomaantha".

Type species. Pseudolomaantha thailandica J.Y. Zhang, Y.Z. Lu & K.D. Hyde Description. Saprobic on dead stems of bamboo in terrestrial habitats. Sexual morph Undetermined. Asexual morph Colonies on natural substrate, effuse, scattered, hairy, dark brown, glistening. Mycelium partly immersed, composed of brown hyphae. Conidiophores macronematous, mononematous, cylindrical, straight or slightly flexuous, septate, dark brown to pale brown. Conidiogenous cells integrated, terminal, holoblastic, monoblastic, cylindrical, brown or pale brown at the apex. Conidia acrogenous, solitary, rostrate, tapering to the round apex, truncate at base, straight or slightly curved, septate, with distoseptate, pale brown to dark brown; with a gold and glistening sheath near the apex.

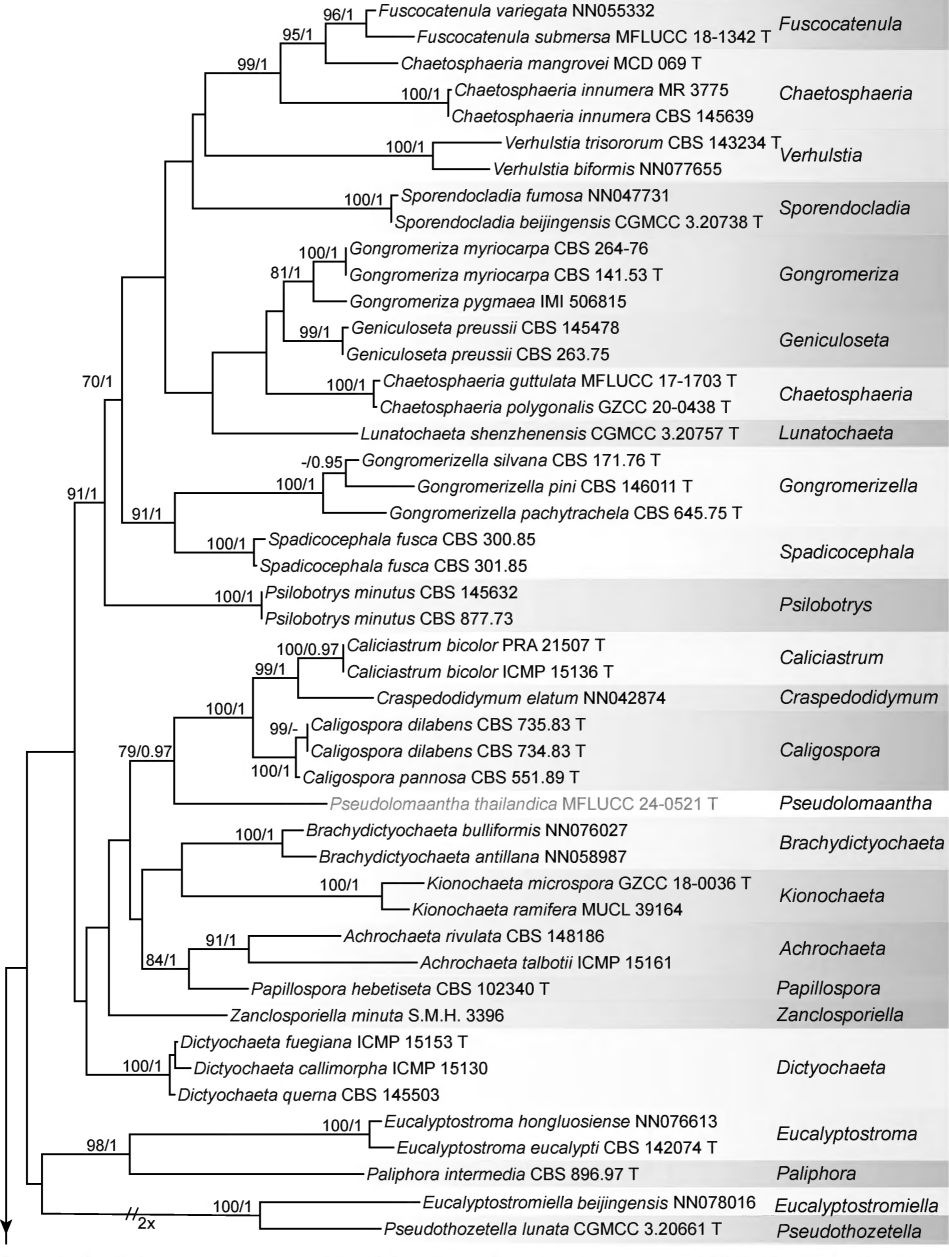
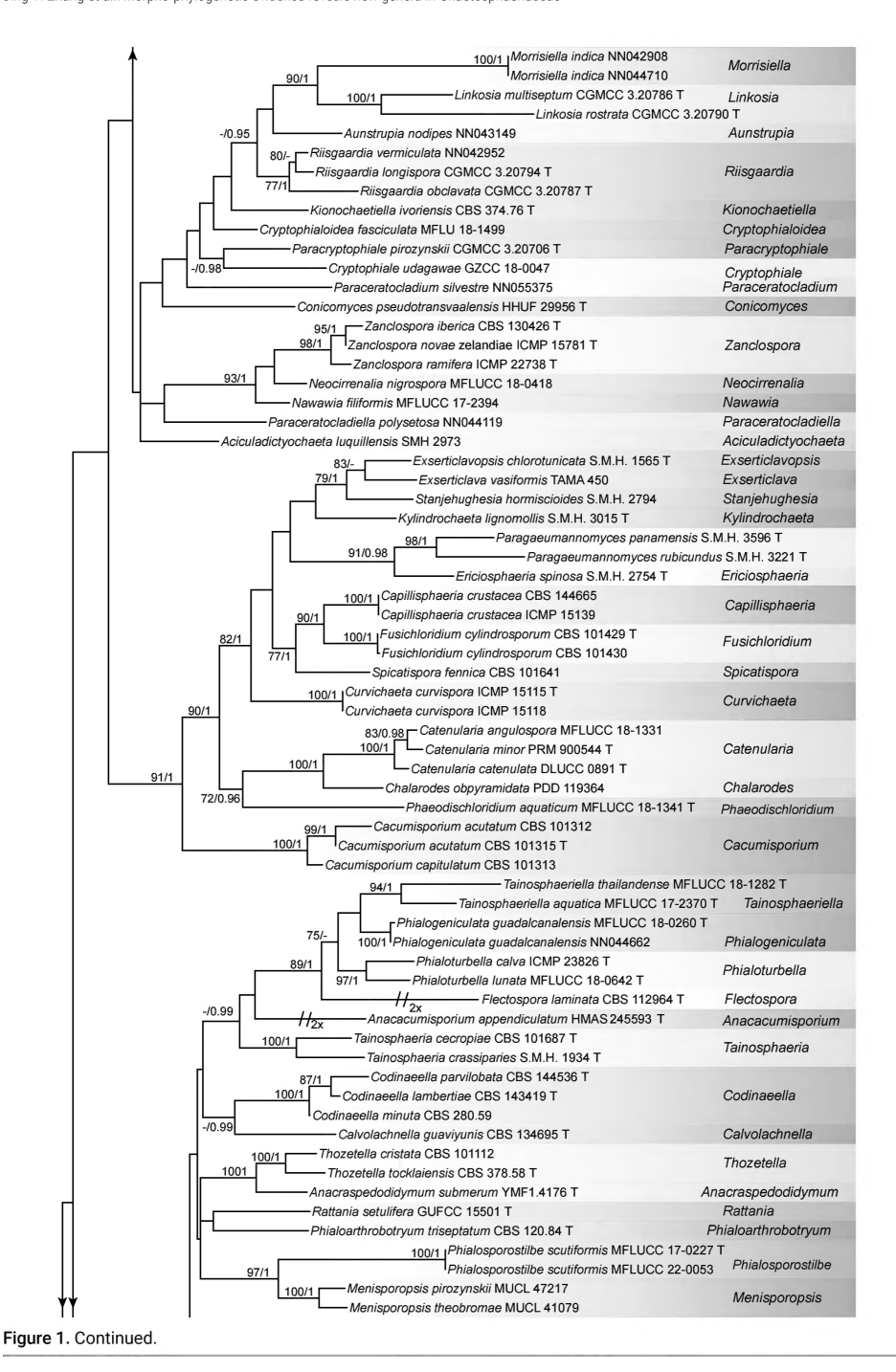


Figure 1. The phylogenetic tree generated from ML analysis is based on a concatenated LSU-ITS dataset for the Chaeto-sphaeriaceae family. Bootstrap support values for ML equal to or greater than 75% and Bayesian posterior probabilities (PPs) equal to or greater than 0.95 were indicated above or below the nodes as ML/PP. *Leptosporella arengae* (MFLUCC 15–0330) and *L. bambusae* (MFLUCC 12–0846) were selected as the outgroup taxa. The newly obtained sequences are indicated in red.



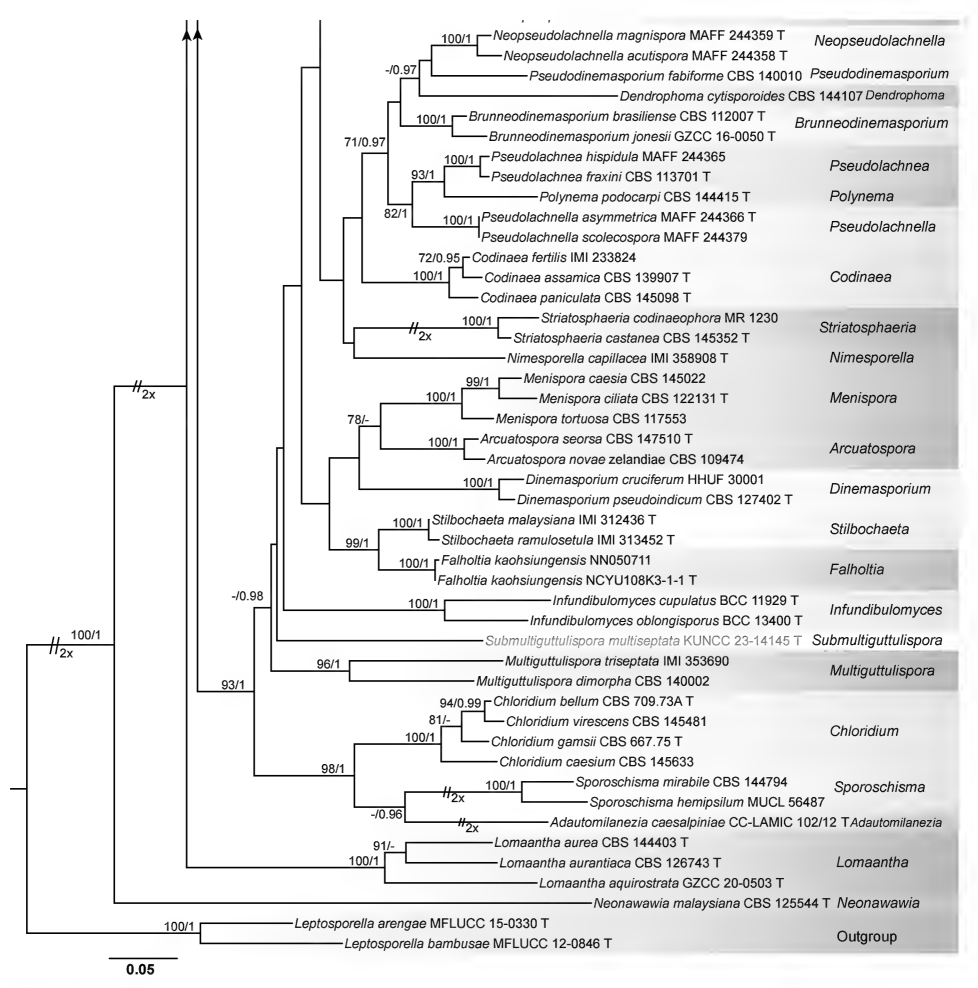


Figure 1. Continued.

Pseudolomaantha thailandica J.Y. Zhang, Y.Z. Lu & K.D. Hyde, sp. nov.

Index Fungorum: IF903138

Facesoffungi Number: FoF16984

Fig. 2

Etymology. The name refers to the country "Thailand" from where the holotype was collected.

Holotype. MFLU 24-0394.

Description. Saprobic on dead stems of bamboo in a terrestrial habitat. Sexual morph Undetermined. Asexual morph Hyphomycetous. Colonies on natural substrate superficial, effuse, scattered, hairy, dark brown, with gold glistening on the apex of conidia. Mycelium partly immersed, partly super-

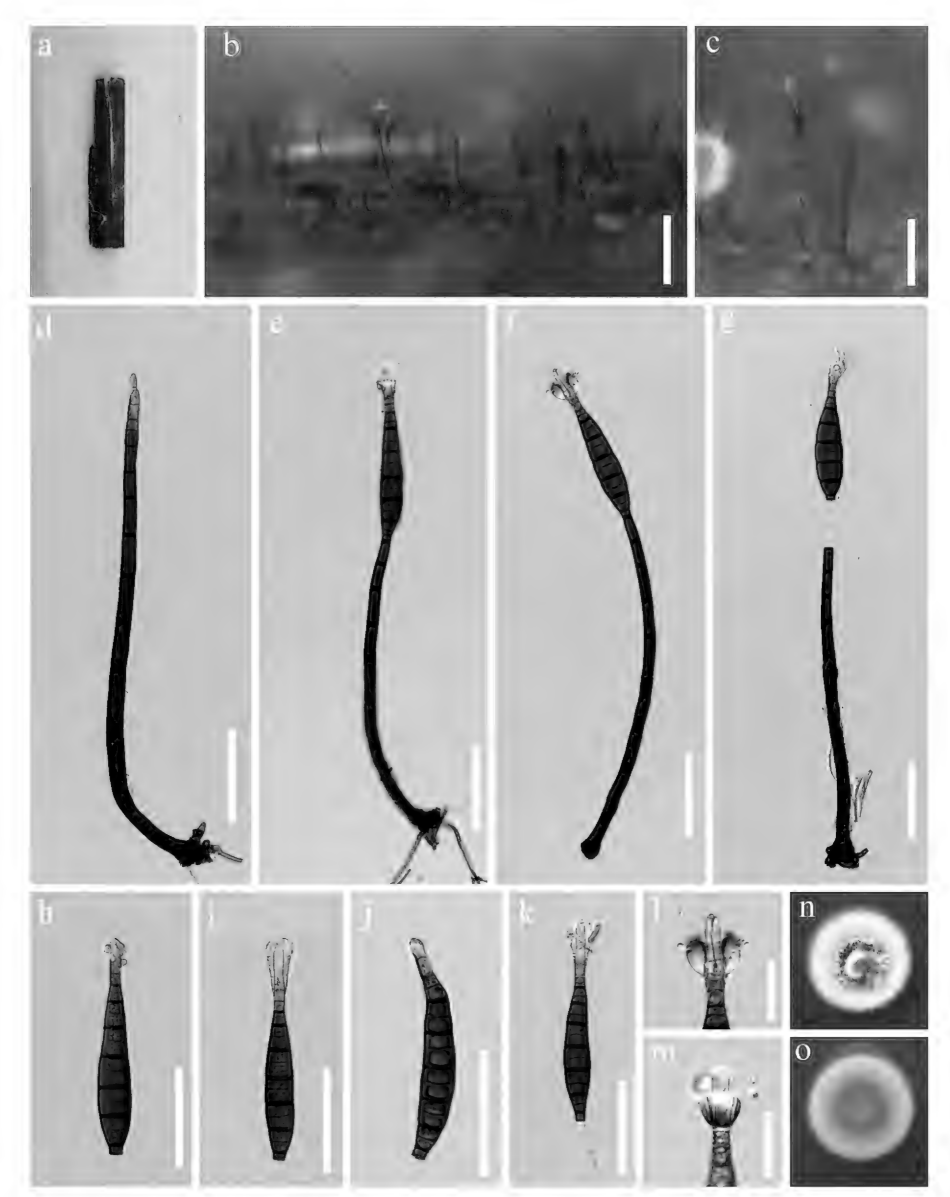


Figure 2. Pseudolomaantha thailandica (MFLU 24–0394, holotype) **a** the host substrate **b**, **c** colonies on the host substrate **d**–**g** conidiophores with conidiogenous cells **h**–**k** conidia **l**, **m** conidial appendage **n**, **o** pure culture from front and reverse. Scale bars: 200 μ m (**b**); 100 μ m (**c**); 50 μ m (**d**–**k**); 20 μ m (**l**, **m**).

ficial, composed of septate, mostly unbranched, smooth, brown hyphae. **Conidiophores** $176-275 \times 6-9(-11) \, \mu m$ ($\bar{x} = 219.6 \times 7.5 \, \mu m$, n = 20), macronematous, mononematous, solitary, cylindrical, straight or slightly flexuous, septate, black at the base, paler to light brown or brown towards the apex.

Conidiogenous cells $12-22 \times 5.5-7 \ \mu m \ (\bar{x}=16.4 \times 6.1 \ \mu m, n=20)$, integrated, terminal, holoblastic, monoblastic, cylindrical, brown or pale brown at the apex. Conidia $(92.5-)95-112.5 \times 12.5-15.5 \ \mu m \ (\bar{x}=105.8 \times 13.8 \ \mu m, n=25)$, acrogenous, solitary, dry, pyriform to obclavate, rostrate, tapering to the round apex, truncate at base, basal cell conical-truncate, straight or slightly curved, up to 12-septate, with distoseptate, not constricted or slightly constricted at septum, guttulate, brown, two upper cells subhyaline to hyaline, with gold and glistening appendages around the apex of the conidia.

Culture characteristics. Conidia germinating on WA within 15 h and germ tube produced from the ends of conidia. Colonies growing on PDA, reaching 22–26 mm in 20 days at 26 °C, circular, edge entire, umbonate with a knobby protuberance, white from above; zonate, yellowish orange in the center, grayish olive to yellowish towards to margin from below.

Material examined. THAILAND • Chiangmai Province, Mushroom Research Center (MRC), on dead stems of bamboo, 11 September 2020, H.W. Shen, Y205-1 (MFLU 24–0394, *holotype*), ex-type living culture, MFLUCC 24–0521.

Notes. BLAST results for the ITS and LSU sequence data of Pseudolomaantha thailandica show 88.39% similarities with Caligospora dilabens (CBS 735.83) and 97.81% similarities with *Craspedodidymum elatum* (NN042874), respectively. Phylogenetic analysis shows that Pseudolomaantha thailandica forms a distinct lineage basal to Caliciastrum, Caligospora, and Craspedodidymum with statistical support (79% ML/0.97 PP, Fig. 1). Members of Caliciastrum, Caligospora, and Craspedodidymum are characterized by phialidic conidiogenous cells with open, vase-shaped collarettes, and brown or hyaline conidia. In contrast, our new species has a sporidesmium-like asexual morph with non-phialidic conidiogenous cells (Figueroa et al. 2018; Wu and Diao 2022; Réblová and Nekvindová 2023). Morphologically, Pseudolomaantha resembles Lomaantha in having macronematous, mononematous conidiophores, integrated holoblastic conidiogenous cells, and acrogenous, obclavate, rostrate, distoseptate, pale brown to brown conidia (Wu and Zhuang 2005; Wu and Diao 2022; Réblová and Nekvindová 2023). However, the two genera are phylogenetically distinct. Additionally, Lomaantha species have conidiogenous cells that are determinate or extend percurrently a few times, as well as conidia that lack or bear filiform, extended, simple or branched apical appendages and distinct septal pores (Wu and Zhuang 2005; Wu and Diao 2022; Réblová and Nekvindová 2023). In contrast, Pseudolomaantha has determinate conidiogenous cells, conidia with golden, glistening appendages at the conidial apex, and lack distinct pores in the distosepta. Based on the combination of morphological and phylogenetic evidence, Pseudolomaantha is introduced as a new genus to accommodate P. thailandica within Chaetosphaeriaceae.

Submultiguttulispora J.Y. Zhang, Y.Z. Lu & K.D. Hyde, gen. nov.

Index Fungorum: IF903141

Facesoffungi Number: FoF16985

Etymology. The name refers to the new genus's close affinity with the genus "Multiguttulispora".

Type species. Submultiguttulispora multiseptatum J.Y. Zhang, Y.Z. Lu & K.D. Hyde.

Description. Saprobic on dead wood. Sexual morph Undetermined. Asexual morph Colonies on natural substrate, effuse, single, or gregarious, brown to black. Mycelium partly immersed, composed of brown hyphae. Conidiophores macronematous, mononematous, single or in small groups, septate, dark brown at the base becoming light brown towards the apex. Conidiogenous cells integrated, mono- to polyphialidic, terminal to lateral, with funnel-shaped collarettes, cylindrical to cylindrical-lageniform, brown to pale brown to subhyaline towards the apex. Conidia acropleurogenous, septate, pale brown to olive green to brown, with subhyaline cells at both ends of the conidia, fusiform, or ellipsoidal-fusiform, with a filiform appendage at each end.

Submultiguttulispora multiseptata J.Y. Zhang, K.D. Hyde & Y.Z. Lu, sp. nov.

Index Fungorum: IF903139

Facesoffungi Number: FoF16986

Fig. 3

Etymology. The name refers to the multi-septate conidia of the new species. **Holotype.** HKAS 129868.

Description. Saprobic on a dead wood log by a stream. Sexual morph undetermined. Asexual morph Hyphomycetous. Colonies on natural substrate superficial, effuse, single, or gregarious, arise in groups from knots of hyphal cells, brown to black. *Mycelium* partly superficial, partly immersed, composed of septate, pale brown to brown, smooth-walled hyphae. Conidiophores 285-385(-533) μ m long × 5–7 μ m wide at the base (\bar{x} = 341 × 6 μ m, n = 15), macronematous, mononematous, single or clustered in groups, erect, straight or flexible, unbranched, septate, smooth, guttulate, dark brown or black at the base, becoming pale brown towards the apex. Conidiogenous cells 64.5-100 × 4.3-6.1 μ m (\bar{x} = 80.2 × 5.2 μ m, n = 15), mono- to polyphialidic, with discrete, terminal to lateral phialides, integrated, terminal, with lateral openings formed by successive sympodial elongation, cylindrical to cylindrical-lageniform, with funnel-shaped collarettes, smooth-walled, guttulate, brown at the base and becoming pale brown to subhyaline towards the apex. Conidia $33-40 \times 7.5-9 \mu m$ $(\bar{x} = 36.6 \times 8.3 \,\mu\text{m}, \, n = 20)$, acropleurogenous, 5(-6)-septate, not constricted at the septum, pale brown to olive green to brown, with subhyaline cells at both ends, straight, sometimes slightly curved, occasionally guttulate, fusiform, or ellipsoidal-fusiform, with a filiform, short and hyaline appendage at each end.

Culture characteristics. Conidia germinating on WA within 15 h and germ tube produced from conidia. Colonies growing on PDA, reaching 35–40 mm diameter in 15 days at 26 °C, circular with slightly irregular edge, flat with a protuberance in the center, dry, velvety, zonate, tephrosiousto to grey from center to margin; dark brown or black from below.

Material examined. CHINA • Hainan Province, Wuzhishan City, Wuzhishan Tropical Rainforest Scenic Area, on a dead wood log by a stream, 15 August 2021, J.Y. Zhang, WZ44-1 (HKAS 129868, holotype; GZAAS 23-0763, isotype); ex-type living cultures, KUNCC 23-14145.

Notes. Based on a BLASTn search in GenBank, the ITS and LSU sequences of our new collection show 91.92% and 95.35% similarity to *Phialogeniculata* guadalcanalensis (NN044662) and *Multiguttulispora triseptata* (IMI 353690),

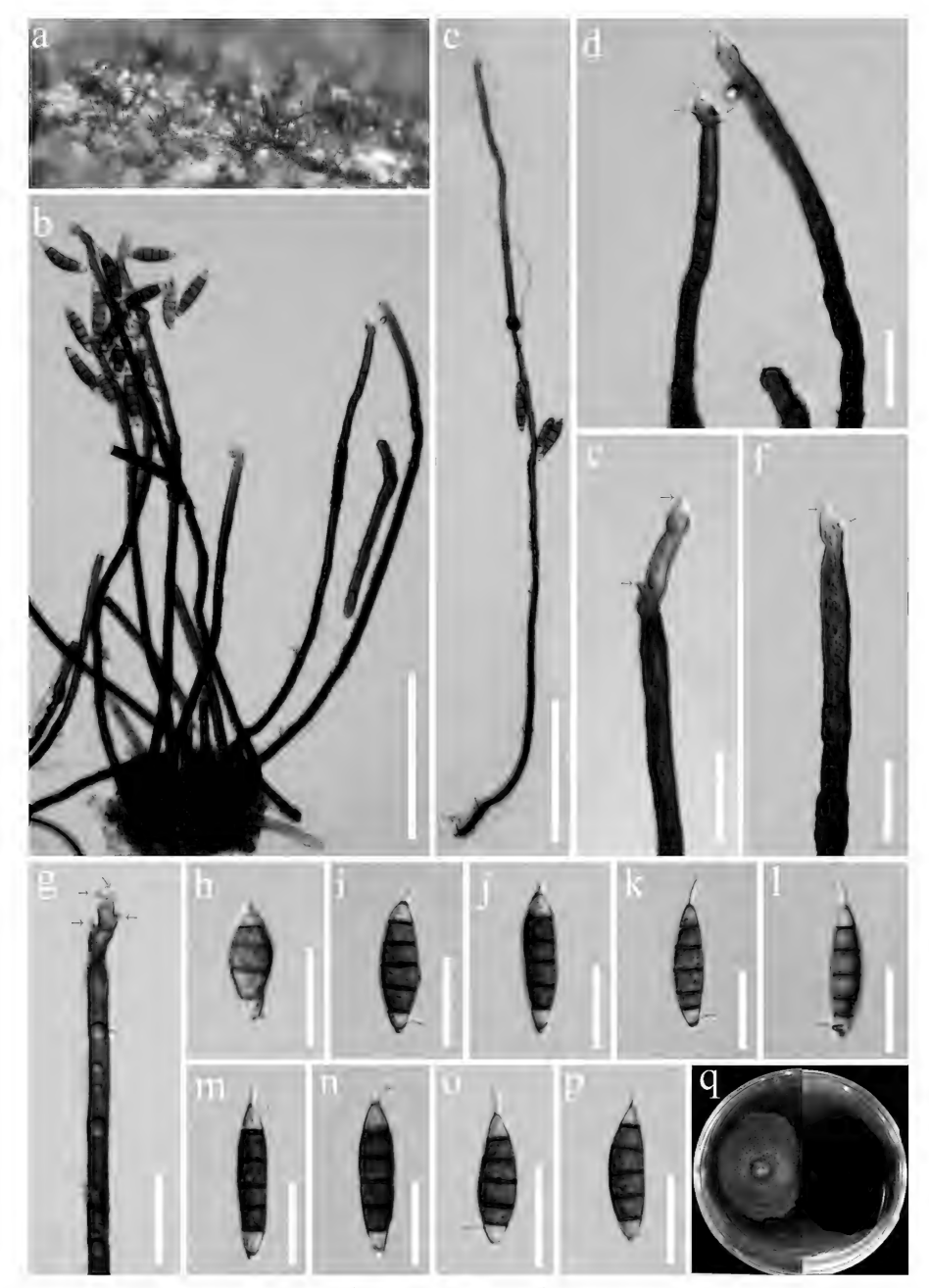


Figure 3. Submultiguttulispora multiseptata (HKAS 129868, holotype) **a** colonies on the host substratum **b**, **c** conidiophores **d**–**g** conidiogenous cells (arrows showing conidiogenous loci) **h**–**p** conidia **q** pure culture from front and reverse. Scale bars: 100 μ m (**b**, **c**); 20 μ m (**d**–**p**).

respectively. The phylogenetic tree indicates that our new isolate forms a distinct lineage closely related to *Multiguttulispora*, without statistical support. This lack of support may be attributed to the absence of molecular sequences of many close phylogenetic relatives, which remain undiscovered (Hyde et al. 2024c). *Submultiguttulispora* shares similarities with *Multiguttulispora* in the absence of setae and the presence of macronematous conidiophores with polyphialidic conidiogenous cells that exhibit sympodial extension. Both genera produce septate conidia with a filiform, hyaline appendage at each end. However, *Submultiguttulispora* is distinguished from *Multiguttulispora* by its fusiform or ellipsoidal-fusiform, dematiaceous conidia, whereas the conidia of *Multiguttulispora* are cylindrical, oblong, and hyaline. Based on these morphological and phylogenetic differences, a new genus, *Submultiguttulispora*, is introduced to accommodate our new isolate, *S. multiseptata*.

Discussion

In this study, *Pseudolomaantha thailandica* gen. et sp. nov. and *Submultiguttu-lispora multiseptata* gen. et sp. nov. were introduced based on morphological characteristics and phylogenetic analyses. These two species exhibit non-phialidic and phialidic asexual morphs, respectively. The introduction of these new taxa further highlights the richness and diversity of anamorphic chaetosphaeriaceous fungi (Réblová et al. 2021a, b, c, d, e; Wu and Diao 2022).

The characteristics of conidiophores, conidiogenous cells, and conidia are particularly important for delimiting asexual genera in Chaetosphaeriaceae, along with the presence or absence of appendages (Réblová et al. 1999, 2021b, c, d; Lin et al. 2019; Zheng et al. 2020; Wu and Diao 2022). A significant number of anamorphic chaetosphaeriaceous genera produce hyaline or subhyaline conidia in various shapes, often with filiform, hyaline setulae at the ends, as seen in genera like Arcuatospora, Codinaea, and Kinochaeta (Hughes and Kendrick 1968; Réblová et al. 2020, 2021b, c; Wu and Diao 2022; Hyde et al. 2024b). In contrast, many hyphomycetous genera with dematiaceous conidia lack setulae, such as Catenularia, Phaeodischloridium and Sporoschisma (Goh et al. 1997; Yang et al. 2016; Réblová et al. 2021e; Wu and Diao 2022). Submultiguttulispora multiseptata gen. et sp. nov. resembles other anamorphic chaetosphaeriaceous genera in having phialidic conidiogenous cells and conidia with filiform, hyaline setulae at both ends (Réblová and Gams 2000; Liu et al. 2016; Lin et al. 2019; Wu and Diao 2022). However, it is distinct in its well-developed conidiophores, polyphialidic conidiogenous cells, and pale brown to olive green to brown, septate conidia with hyaline setulae at each end. The latest key to phialidic asexual genera in Chaetosphaeriaceae was provided by Wu and Diao (2022).

Wu and Diao (2022) recognized ten hyphomycetous genera with non-phialidic anamorphs in Chaetosphaeriaceae, viz., Aunstrupia, Ellisembia, Falholtia, Linkosia, Lomaantha, Morrisiella, Paliphora, Riisgaardia, Stanjehughesia, and Zanclospora. Subsequently, a new non-phialidic genus, Neocirrenalia, characterized by dark brown or black helicoid conidia, was added to this family (Meyers and Moore 1960; Somrithipol et al. 2002; Yang et al. 2023). Recently, Delgado et al. (2024) reclassified Ellisembia into Sporidesmiaceae (Sordariomycetes) based on analyses of a newly collected type species, E. coronata, and expanded and emended Lomaantha to include related ellisembia-like taxa

within a monophyletic lineage in Chaetosphaeriaceae. Currently, Chaetosphaeriaceae comprises 10 non-phialidic hyphomycetous genera. Most of these genera are sporidesmium-like, with exceptions such as *Neocirrenalia* (a helicosporous genus) and *Paliphora*, which is characterized by setiform conidiophores, polytretic conidiogenous cells, and subfusiform to subacerose, hyaline conidia (Gusmão et al. 2008; Shenoy et al. 2010; Goh et al. 2014; Malosso et al. 2017; Wu and Diao 2022; Yang et al. 2023; Ma et al. 2024). In this study, we introduced a new genus, *Pseudolomaantha*, which also exhibits a sporidesmium-like asexual morph, characterized by well-developed, solitary or clustered conidiophores and pyriform to obclavate conidia with a glistening gold appendage around the apex, but is phylogenetically distinct. A key to hyphomycetous genera with non-phialidic anamorphs is provided herein.

Key to hyphomycetous genera with non-phialidic anamorphs

1	Sporidesmium-like genera2
_	Not sporidesmium-like genera3
2	Conidiophores absent (reduced to conidiogenous cells), or solitary or in a small group
_	Conidiophores in synnemata5
3	Conidiogenous cells polytretic; conidia hyaline, subfusiform to subacerose
_	Conidiogenous cells monoblastic; conidia black, helicoid Neocirrenalia
4	Conidia with appendage at the apex6
_	Conidia without appendage7
5	Conidia euseptate
_	Conidia distoseptate
6	Conidiophores absent or well-developed, conidia cylindrical, obclavate or
	narrowly fusiform, often with filamentous, hyaline apical appendages and
	typically bearing distinct pores in the distosepta
_	Conidiophores well-developed, conidia pyriform to obclavate, with a gold
	and glistening appendage around the apex, and distinct pores are not ob-
	served
7	Synanamorph of Zanclospora with phialides Zanclospora
-	Not synanamorph of Zanclospora8
8	Conidiophores absent; conidia euseptate9
-	Conidiophores absent, conidia distoseptate
9	Conidia obclavate, obclavate-rostrate, subcylindrical
-	Conidia cylindrical, clavate, or obclavate

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Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

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Author contributions

Morphological data, photo plates, and phylogenetic analyses were completed by Jing-Yi Zhang. The original draft was written by Jing-Yi Zhang, and Kevin D. Hyde, Jian Ma, Na Wu, Fatimah Al-Otibi & Yong-Zhong Lu revised the paper. Financial support was provided by Li-Juan Zhang and Yong-Zhong Lu.

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Data availability

All of the data that support the findings of this study are available in the main text.

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